Christoph Bannwarth

List of Publications by Year in Descending Order

Source: https://exaly.com/author-pdf/8732922/christoph-bannwarth-publications-by-year.pdf

Version: 2024-04-20

This document has been generated based on the publications and citations recorded by exaly.com. For the latest version of this publication list, visit the link given above.

The third column is the impact factor (IF) of the journal, and the fourth column is the number of citations of the article.

55
papers

4,898
citations

25
h-index

67
g-index

67
ext. papers

25
citations

8.7
ext. citations

8.7
ext. citations

8.7
ext. citations

L-index

#	Paper	IF	Citations
55	Chiral photochemistry of achiral molecules <i>Nature Communications</i> , 2022 , 13, 2091	17.4	1
54	Predictions of Pre-edge Features in Time-Resolved Near-Edge X-ray Absorption Fine Structure Spectroscopy from Hole-Hole Tamm-Dancoff-Approximated Density Functional Theory. <i>Journal of Chemical Theory and Computation</i> , 2021 , 17, 7120-7133	6.4	О
53	Hot exciplexes in U-shaped TADF molecules with emission from locally excited states. <i>Nature Communications</i> , 2021 , 12, 6179	17.4	7
52	Extended tight-binding quantum chemistry methods. Wiley Interdisciplinary Reviews: Computational Molecular Science, 2021 , 11, e1493	7.9	149
51	TeraChem: A graphical processing unit-accelerated electronic structure package for large-scale ab initio molecular dynamics. <i>Wiley Interdisciplinary Reviews: Computational Molecular Science</i> , 2021 , 11, e1494	7.9	49
50	Cover Image, Volume 11, Issue 2. <i>Wiley Interdisciplinary Reviews: Computational Molecular Science</i> , 2021 , 11, e1523	7.9	3
49	Photochemical Deracemization of Primary Allene Amides by Triplet Energy Transfer: A Combined Synthetic and Theoretical Study. <i>Journal of the American Chemical Society</i> , 2021 , 143, 11209-11217	16.4	12
48	Cycloadditions of Donor-Acceptor Cyclopropanes and -butanes using S=N-Containing Reagents: Access to Cyclic Sulfinamides, Sulfonamides, and Sulfinamidines. <i>Angewandte Chemie - International Edition</i> , 2021 , 60, 25825-25831	16.4	2
47	TeraChem: Accelerating electronic structure and ab initio molecular dynamics with graphical processing units. <i>Journal of Chemical Physics</i> , 2020 , 152, 224110	3.9	40
46	Hole-hole Tamm-Dancoff-approximated density functional theory: A highly efficient electronic structure method incorporating dynamic and static correlation. <i>Journal of Chemical Physics</i> , 2020 , 153, 024110	3.9	15
45	Nonadiabatic Dynamics Simulation of the Wavelength-Dependent Photochemistry of Azobenzene Excited to the nB and B Excited States. <i>Journal of the American Chemical Society</i> , 2020 , 142, 20680-20690	o ^{16.4}	20
44	Nonadiabatic Molecular Dynamics with Hole-Hole Tamm-Dancoff Approximated Density Functional Theory. <i>Journal of Chemical Theory and Computation</i> , 2020 , 16, 5499-5511	6.4	12
43	A Simplified Spin-Flip Time-Dependent Density Functional Theory Approach for the Electronic Excitation Spectra of Very Large Diradicals. <i>Journal of Physical Chemistry A</i> , 2019 , 123, 5815-5825	2.8	9
42	A generally applicable atomic-charge dependent London dispersion correction. <i>Journal of Chemical Physics</i> , 2019 , 150, 154122	3.9	300
41	GFN2-xTB-An Accurate and Broadly Parametrized Self-Consistent Tight-Binding Quantum Chemical Method with Multipole Electrostatics and Density-Dependent Dispersion Contributions. <i>Journal of Chemical Theory and Computation</i> , 2019 , 15, 1652-1671	6.4	717
40	Extension of the element parameter set for ultra-fast excitation spectra calculation (sTDA-xTB). <i>Molecular Physics</i> , 2019 , 117, 1104-1116	1.7	3
39	B97-3c: A revised low-cost variant of the B97-D density functional method. <i>Journal of Chemical Physics</i> , 2018 , 148, 064104	3.9	221

(2016-2018)

38	Catalytic deracemization of chiral allenes by sensitized excitation with visible light. <i>Nature</i> , 2018 , 564, 240-243	50.4	95
37	Benzimidazolylquinoxalines: novel fluorophores with tuneable sensitivity to solvent effects. <i>Physical Chemistry Chemical Physics</i> , 2017 , 19, 6095-6104	3.6	10
36	A Robust and Accurate Tight-Binding Quantum Chemical Method for Structures, Vibrational Frequencies, and Noncovalent Interactions of Large Molecular Systems Parametrized for All spd-Block Elements (Z = 1-86). <i>Journal of Chemical Theory and Computation</i> , 2017 , 13, 1989-2009	6.4	592
35	Ein achtkerniger metallosupramolekularer WEFel mit Spin-Crossover-Eigenschaften. <i>Angewandte Chemie</i> , 2017 , 129, 5012-5017	3.6	13
34	Frontispiece: An Octanuclear Metallosupramolecular Cage Designed To Exhibit Spin-Crossover Behavior. <i>Angewandte Chemie - International Edition</i> , 2017 , 56,	16.4	1
33	Pyridyl Containing 1,5-Diaza-3,7-diphosphacyclooctanes as Bridging Ligands for Dinuclear Copper(I) Complexes. <i>Zeitschrift Fur Anorganische Und Allgemeine Chemie</i> , 2017 , 643, 895-902	1.3	11
32	An Octanuclear Metallosupramolecular Cage Designed To Exhibit Spin-Crossover Behavior. <i>Angewandte Chemie - International Edition</i> , 2017 , 56, 4930-4935	16.4	59
31	Vollautomatisierte quantenchemische Berechnung von Spin-Spin- gekoppelten magnetischen Kernspinresonanzspektren. <i>Angewandte Chemie</i> , 2017 , 129, 14958-14964	3.6	25
30	Fully Automated Quantum-Chemistry-Based Computation of Spin-Spin-Coupled Nuclear Magnetic Resonance Spectra. <i>Angewandte Chemie - International Edition</i> , 2017 , 56, 14763-14769	16.4	107
29	A general intermolecular force field based on tight-binding quantum chemical calculations. <i>Journal of Chemical Physics</i> , 2017 , 147, 161708	3.9	36
28	Extension of the D3 dispersion coefficient model. <i>Journal of Chemical Physics</i> , 2017 , 147, 034112	3.9	293
27	Biomolecular Structure Information from High-Speed Quantum Mechanical Electronic Spectra Calculation. <i>Journal of the American Chemical Society</i> , 2017 , 139, 11682-11685	16.4	24
26	Diastereoselective Self-Assembly of a Neutral Dinuclear Double-Stranded Zinc(II) Helicate via Narcissistic Self-Sorting. <i>Chemistry - A European Journal</i> , 2017 , 23, 12380-12386	4.8	16
25	Indirect synthesis of a pair of formal methane activation products at a phosphane/borane frustrated Lewis pair. <i>Dalton Transactions</i> , 2016 , 45, 19230-19233	4.3	7
24	Effect of Conjugation Pathway in Metal-Free Room-Temperature Dual Singlet-Triplet Emitters for Organic Light-Emitting Diodes. <i>Journal of Physical Chemistry Letters</i> , 2016 , 7, 4802-4808	6.4	30
23	Recent research directions in Fribourg: nuclear dynamics in resonances revealed by 2-dimensional EEL spectra, electron collisions with ionic liquids and electronic excitation of pyrimidine. <i>European Physical Journal D</i> , 2016 , 70, 1	1.3	6
22	Electronic Circular Dichroism of [16]Helicene With Simplified TD-DFT: Beyond the Single Structure Approach. <i>Chirality</i> , 2016 , 28, 365-9	2.1	24
21	Non-covalent Stabilization in Transition Metal Coordination and Organometallic Complexes 2016 , 115-1	43	4

20	Ultra-fast computation of electronic spectra for large systems by tight-binding based simplified Tamm-Dancoff approximation (sTDA-xTB). <i>Journal of Chemical Physics</i> , 2016 , 145, 054103	3.9	87
19	Dispersion-Corrected Mean-Field Electronic Structure Methods. <i>Chemical Reviews</i> , 2016 , 116, 5105-54	68.1	738
18	Synthesis and Comprehensive Structural and Chiroptical Characterization of Enones Derived from (-)-ESantonin by Experiment and Theory. <i>Journal of Organic Chemistry</i> , 2016 , 81, 4588-600	4.2	10
17	Direct synthesis of a geminal zwitterionic phosphonium/hydridoborate systemdeveloping an alternative tool for generating frustrated Lewis pair hydrogen activation systems. <i>Organic and Biomolecular Chemistry</i> , 2015 , 13, 5783-92	3.9	22
16	Electronic circular dichroism of highly conjugated Bystems: breakdown of the Tamm-Dancoff/configuration interaction singles approximation. <i>Journal of Physical Chemistry A</i> , 2015 , 119, 3653-62	2.8	15
15	Consistent structures and interactions by density functional theory with small atomic orbital basis sets. <i>Journal of Chemical Physics</i> , 2015 , 143, 054107	3.9	404
14	Enamine/butadienylborane cycloaddition in the frustrated Lewis pair regime. <i>Organic and Biomolecular Chemistry</i> , 2015 , 13, 10477-86	3.9	8
13	Enantiotopos-selektive CH-Oxygenierung mit einem supramolekularen Ruthenium-Katalysator. <i>Angewandte Chemie</i> , 2015 , 127, 701-705	3.6	23
12	Enantiotopos-selective C-H oxygenation catalyzed by a supramolecular ruthenium complex. <i>Angewandte Chemie - International Edition</i> , 2015 , 54, 691-5	16.4	46
11	Free electrons and ionic liquids: study of excited states by means of electron-energy loss spectroscopy and the density functional theory multireference configuration interaction method. <i>Physical Chemistry Chemical Physics</i> , 2015 , 17, 15771-80	3.6	12
10	Synthesis, Chiral Resolution, and Absolute Configuration of Functionalized Trgers Base Derivatives: Part III. <i>Synthesis</i> , 2015 , 47, 3118-3132	2.9	8
9	The Association of Two Brustrated(Lewis Pairs by State-of-the-Art Quantum Chemical Methods. <i>Israel Journal of Chemistry</i> , 2015 , 55, 235-242	3.4	20
8	The thermochemistry of london dispersion-driven transition metal reactions: getting the 'right answer for the right reason'. <i>ChemistryOpen</i> , 2014 , 3, 177-89	2.3	70
7	A simplified time-dependent density functional theory approach for electronic ultraviolet and circular dichroism spectra of very large molecules. <i>Computational and Theoretical Chemistry</i> , 2014 , 1040-1041, 45-53	2	138
6	Enantiomerenreine [M6L12]- oder [M12L24]-Polyeder aus flexiblen Bis(pyridin)-Liganden. <i>Angewandte Chemie</i> , 2014 , 126, 1719-1724	3.6	32
5	Enantiomerically pure [M(6)L(12)] or [M(12)L(24)] polyhedra from flexible bis(pyridine) ligands. <i>Angewandte Chemie - International Edition</i> , 2014 , 53, 1693-8	16.4	86
4	Combinations of ethers and B(C6F5)3 function as hydrogenation catalysts. <i>Angewandte Chemie - International Edition</i> , 2013 , 52, 7492-5	16.4	160
3	Combinations of Ethers and B(C6F5)3 Function as Hydrogenation Catalysts. <i>Angewandte Chemie</i> , 2013 , 125, 7640-7643	3.6	64

LIST OF PUBLICATIONS

From attraction to repulsion: anion-linteractions between bromide and fluorinated phenyl groups. Chemical Communications, **2011**, 47, 8542-4

5.8 37

A Generally Applicable Atomic-Charge Dependent London Dispersion Correction Scheme

2